

The stability and melting of aragonite: an experimental and thermodynamic model for carbonated eclogites in the mantle

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1. Introduction

Calcium carbonate is the primary compound of carbon reservoirs in the Earth’s oceanic and continental crust. Biomineralization in the oceans is responsible for accumulation of hundred meters thick sequences of nannofossil oozes and chalks almost entirely composed of pure calcite (e.g., DSDP site 501 and 504 in the East Pacific, and site 212 in the Indian Ocean; Cann et al., 1983, and von der Borch et al., 1974). In the underlying igneous suites, hydrothermal fluids percolate to a depth of several kilometers, triggering the widespread precipitation of calcite and aragonite in vugs and veins (Bonatti et al. 1980; Alt and Teagle, 1999). Again, calcite cements serpentized ultramafics in ophicarbonates breccias, which are produced in the highly tectonized oceanic lithosphere, at slow spreading ridges, and at transform faults in fast spreading oceans. At continental margins, carbonate platforms are almost entirely built by organism fixing calcium carbonate.

Subduction and mantle convection are responsible for high-pressure and high-temperature reprocessing of Ca-carbonates in a diversity of thermo-mechanical scenarios, accounting for CO₂ sink in the frame of the global, long-term carbon cycle. Aragonite has been regarded since long time

as a marker of relatively low temperature - high pressure metamorphism, and it has been widely used in reference geobarometers for so-called ultra-high pressure metamorphic rocks (Hermann et al., 2016). Nonetheless, as Ca-carbonates are stable over a very wide range of temperatures (Suito et al., 2001; Li et al., 2017; Müller et al., 2017; Shatskiy et al., 2018), up to the average mantle adiabat beneath ridges and ocean islands (Fig. 1), a large range of carbonated eclogites and pyroxenites may contain aragonite at mantle conditions, as revealed by several experimental studies (Hammouda, 2003; Kiseeva et al., 2013; Grassi and Schmidt, 2011; Brey et al., 2015). Evidence of CaCO_3 -bearing eclogitic assemblages have been also provided by the occurrence of inclusions of carbonates in the diamonds from Juina (Brazil), possibly originated from the lower part of the transition zone (Brenker et al., 2007), and of aragonite as inclusions in olivine phenocrysts from leucitite lava flows at Calatrava, Spain (Humphreys et al., 2010).

The geochemistry of carbonatites (Woolley and Kjarsgaard, 2008) and kimberlites (Becker and Le Roex, 2006) points to the importance of components CaO and CO_2 for describing their diversity and magmatic evolution. Carbonatites are thought to be among the major metasomatic agents in the sublithospheric mantle due to their low density, low viscosity and high reactivity (Green and Wallace, 1988). Whether calcite or aragonite occur on the liquidus surface it may affect the fractionation of trace elements as aragonite was experimentally found to preferentially partition Sr with respect to calcite (Carlson, 1980), and intermediate REE with respect to Fe-Mg carbonate (Brey et al., 2015).

Thermodynamic properties of CaCO_3 polymorphs, and of liquid CaCO_3 , are barely known (Salje and Viswanathan, 1976; Redfern et al., 1989; Antao et al., 2009; Liu and Lange, 2003; O'Leary et al., 2015; Vuilleumier et al., 2014). Moreover, the volume equation of state of calcite V is substantially unknown at high temperature. Notably, the phase transition at high temperature from CaCO_3 -polymorph calcite V to aragonite shows large discrepancies among the existing results (Fig. 1), and the behaviour of calcite V has been previously modelled extrapolating low-temperature

51 experimental data. Thermodynamic properties for a CaCO_3 component in liquid (hereafter CaCO_3L)
52 is crucial for developing multi-component solution models suitable for predicting melting relations
53 in carbonated mantle rocks, profiting of properties already available for FeCO_3L (Kang et al., 2015)
54 and MgCO_3L (Kang et al., 2016).

55 Here we present an experimental study for the system CaCO_3 conducted with a multi-anvil
56 apparatus at 3-6 GPa and 1300-1750 °C. In order to constrain the high temperature volume equation
57 of state of calcite and aragonite, we additionally present in-situ X-ray diffraction experiments on
58 calcite up to 1000 °C. Thermodynamic properties of aragonite, calcite V, and liquid CaCO_3 are
59 derived by least-square fitting of experimental results and are used to calculate phase relations in
60 carbonated eclogite composition where saturation in calcium carbonate is predicted over a wide
61 range of pressures and temperatures, up to the *solidus*.

62 **2. Experimental background on the system CaCO_3**

63 Calcium carbonate (CaCO_3) is found in nature in three different structural forms: calcite (R-
64 3c), aragonite (Pmcn), and vaterite (P63/mmc), although the stability field of vaterite is not well
65 known (Maruyama et al., 2017). Aragonite generally represents the high-pressure and relatively
66 low-temperature stable structure of CaCO_3 , but it is commonly found as a metastable phase within
67 the stability field of calcite, even at ambient conditions, due to its growth kinetics. At higher
68 pressure, aragonite will transform to a post-aragonite phase at about 40 GPa (Merlini et al., 2016;
69 Ono et al., 2007; Palaich et al., 2016).

70 Calcite has been reported with six different modification structures (calcite I, II, III, IV, V, and
71 VI) over a wide range of pressure and temperature (Bagdassarov and Slutskii, 2003; Litasov et al.,
72 2017; Mirwald, 1976; Redfern et al., 1989; Merlini et al., 2012). At room temperature, calcite I
73 transforms into calcite II (monoclinic structure, P21/c) at 1.7 GPa and then to calcite III (C2) at 2.0
74 GPa (Smyth and Ahrens, 1997), which appears to persist to more than 6 GPa at very low-T

75 conditions (Suito et al., 2001). Calcite V, commonly regarded as disordered calcite, has been found
76 at temperature higher than approximately 966 °C (Bagdassarov and Slutskii, 2003; Mirwald, 1976;
77 Ishizawa et al., 2013), while calcite IV can be considered as an intermediate phase between calcite I
78 to V related to disordering of the CO_3^{2-} groups (Ishizawa et al., 2013), above 700 °C.
79 Transformation from phase IV into V is marked by the change of the space group from $R\bar{3}c$ to $R\bar{3}m$,
80 with the c-length being halved. In Phase V, the oxygen atoms exist with equal probability along the
81 undulated circular orbital around the central carbon.

82 The polymorphic transformation from trigonal calcite to orthorhombic aragonite is the most
83 visible phase transition in the system CaCO_3 . Nonetheless, previous studies aimed at defining the
84 calcite-aragonite boundary yielded contradictory results, especially over the range of temperatures
85 where calcite V is the stable low pressure phase (Fig. 1).

86 Experiments by Irving and Wyllie (1975) up to 3.5 GPa covered a temperature range from 800
87 °C to 1100 °C, and represent the solely synthesis reversals available where mixtures of both calcite
88 and aragonite were used as starting materials. Their experimental data agree fairly well with the
89 phase boundary calculated by Redfern et al. (1989) based on calorimetric measurements, and
90 extrapolation of the calcite-aragonite transition curve to higher temperatures intersects CaCO_3 -
91 melting curve at an invariant point approximately located at 5.5 GPa and ~ 1650 °C (Irving and
92 Wyllie, 1975). However, these results appear to be in poor agreement with DTA investigations on
93 pure calcite by Mirwald (1976), and even contrast with in situ X-ray diffraction studies by Litasov
94 et al. (2017), and Suito et al. (2001), where a disordered calcite was observed up to 6 GPa and 1230
95 °C that remains stable until melting (Fig. 1); the boundary aragonite-calcite suggested by Litasov et
96 al. (2017) progressively increases its dP/dT slope until it becomes pressure insensitive above ~ 10
97 GPa at ~ 1300 °C. In-situ experiments are in substantial contrast with experimental results by Buob
98 (2003), where CaCO_3 was found to maintain the aragonite structure up to 6 GPa and temperatures

99 between 1300-1500 °C, and by Shatskiy et al. (2014), where melting of aragonite was detected
100 between 1600-1700 °C at 6 GPa.

101 The melting curve for CaCO_3 was recently investigated by Li et al. (2017) in the pressure
102 range 3-21 GPa, based on *in situ* conductivity experiments; temperature of melting was detected on
103 the basis of a steep and large rise in the ionic current through the sample during each heating cycle.
104 Pre-melting peaks in the ionic current were attributed to phase transition from aragonite to calcite
105 V, but results from marker experiments and Raman measurements of recovered samples exhibit
106 aragonite in the P-T region where calcite V was supposed to be stable (Fig. 3b in Li et al., 2017).
107 Melting of aragonite was bracketed down to 8 GPa by a falling Pt-sphere technique. Shatskiy et al.
108 (2018) revisited the CaCO_3 - MgCO_3 phase diagram and describe aragonite in equilibrium with melt
109 at 6 GPa, in contrast with previous evidence (i.e. Litasov et al., 2017). A more recent paper,
110 focussing on ultra-high pressure phase diagram of CaCO_3 (Bayarjgal et al., 2018) report in-situ
111 experimental data indicating stability of aragonite at 11 GPa at temperatures higher than the
112 stability line proposed by Litasov et al. (2017), indicating that further experimental work is
113 mandatory to unravel the phase diagram reconstruction even at pressures beyond the range
114 discussed here.

115 Currently available thermodynamic databases of petrological interest (Holland and Powell,
116 1998; update 2002; 2011) predict phase transformation of calcite to aragonite close to the
117 experimental results of Mirwald (1976) at temperatures lower than ~ 1200 °C. It is worth noting
118 that the 2002 update of the database by Holland and Powell (1998) includes a Landau model for
119 aragonite, lately removed in Holland and Powell (2011), as aragonite does not exhibit disordering of
120 CO_3 groups.

121 Synthesis, ex-situ, experiments, mostly cover the low-temperature, low-pressure range,
122 whereas *in-situ* investigations mostly focus on the high-temperature, and high-pressure region of the
123 phase diagram. Whether discrepancies should be partly ascribed to inter-laboratory pressure

124 calibration used is currently unclear; nonetheless, uncertainty on the location of the equilibrium
125 boundary close to the *solidus* is such that further data are mandatory to unravel appropriate
126 parameters for predicting the fate of carbonated eclogites at mantle pressure.

127 **3. Experimental and analytical techniques**

128 The experimental study was conducted in a 1000-ton Walker-type multi-anvil *apparatus* at the
129 Università degli Studi di Milano from 1300 °C to 1750 °C and 3 to 6 GPa. Experiments were all
130 performed in a multi-anvil apparatus, in order to provide the internal consistency of the data, after
131 recalibration of the pressure medium assembly (see Supplementary data A). In order to attain
132 textural equilibrium at run conditions, experiments at 1300-1500 °C were performed with a run
133 duration in the order of one day, whereas experiments at 1650-1750 °C lasted 1-2 hours. The
134 starting material was made of reagent-grade pure CaCO₃ powder (purity larger than 99.95%)
135 purchased from BDH Chemical Ltd, and synthetic aragonite.

136 The powders were permanently stored in a vacuum oven at 110 °C to prevent hydration, which
137 is known to cause a strong depression of CaCO₃ *solidus* temperature (Wyllie and Boettcher, 1969;
138 Müller et al 2017). Bracketing experiments were performed using starting materials containing both
139 reactants and products expected, in order to localize accurately the univariant equilibrium in the P-T
140 space (Pattison, 1994).

141 A 25/17 assembly was adopted to minimize thermal gradient in the run charge, using Cr₂O₃
142 doped MgO-octahedra, and preformed pyrophyllite gaskets. The assembly includes ZrO₂ sleeve,
143 cylindrical graphite furnace, internal MgO spacers, and molybdenum end spacers ensuring electrical
144 contact. Temperature was controlled by Eurotherm controllers within ± 2°C, and monitored by S-
145 type (Pt₁₀₀/Pt₉₀Rh₁₀) axial thermocouples. Typical thermal gradient within the assembly is ± 20 °C
146 across the capsule. Starting material was loaded into a platinum capsule, and then stacked in the
147 central part of the furnace to minimize the uncertainty due to the thermal gradient across the sample

148 chamber. In order to reduce the influx of hydrogen during the experiments, as expected by the Soret
149 effect, and then ensure a nearly-anhydrous run conditions, the capsule was packed with Fe_2O_3
150 powder before being embedded in a MgO sleeve (Liu and O'Neill, 2004).

151 At the end of the experiments, assembly was gradually decompressed overnight after
152 quenching by shutting off the heating power. The averaged quench rate in the interval 500-1500 °C
153 is around 400°C/sec, decreasing exponentially at lower temperatures. Recovered samples were
154 longitudinally embedded in epoxy resins, ground and polished with diamond paste in order to
155 expose a level of the product of synthesis, and finally carbon coated for Scanning Electron
156 Microscope (SEM) analysis. Textural analysis of the phase assemblages was carried out by back-
157 scattered (BSE), and secondary electrons imaging using a JEOL JXA8200 electron microprobe.
158 Run products were also characterized by X-ray diffraction using Oxford X'calibur diffractometer
159 (Mo X-ray source) available at the Department of Earth Sciences, University of Milan, to identify
160 the mineral phases.

161 High temperature diffraction on CaCO_3 powder was performed at MCX beamline at Elettra
162 synchrotron facility, using the high temperature furnace available and the standard experimental
163 setup (Riello et al. 2013; Merlini et al., 2016). In-situ diffraction were collected in the temperature
164 interval 25-1000°C, with sample contained in a quartz capillary connected to a CO_2 gas line at 1.5
165 bar to prevent decarbonation during the experiment (see the supplementary data B for details).

166 **4. Results**

167 A total of 11 successful synthesis experiments have been performed. Run conditions, duration
168 of the experiments, and run products are reported in Table 1. Evidence for textural equilibration at
169 subsolidus conditions comes from the development of polygonal grain boundaries and triple
170 junctions between solid phases. Because CaCO_3 liquid phase does not quench to a glass, due to its
171 low viscosity, melting in the system was inferred by the crystal morphologies that are

representatives of a textural disequilibrium, such as fibrous, elongated, or feather-like textures (Fig. 2) (Donaldson, 1976; Paterson, 1958).

The identification of CaCO_3 structural polymorph was based on X-ray powder diffraction (see supplementary data B). Since phase transformations from calcite I to IV, and V are unquenchable, the ex-situ characterization of high pressure experiments provide evidence for calcite I symmetry at ambient conditions for run products at 3.3 and 3.8 GPa, 1300 °C, and 3.8 GPa 1500 °C (Fig. 3). In-situ high-temperature experiments at ambient pressure show the peculiar thermal behaviour of calcite. A negative expansion of the a axis characterizes the structural evolution below approximately 600 °C. At this temperature the calcite to CaCO_3 -IV transition is traditionally located. Whether CaCO_3 -IV is a different polymorph of calcite or it represent a transitional behaviour of calcite is matter of debate. It is noticeable that the refinement of oxygen occupancy (see figure B4, supplementary data B) using a two carbonate group model, decrease starting at 600 °C and reach the 0.5 value of the fully disordered phase (CaCO_3 -V) around 1000 °C.

Aragonite was detected at 4.2 GPa and 1300 °C, and remains stable up to 1650 °C at 4.7 GPa (Table 1, Fig. 3). It is extremely unlikely that aragonite occurrence in reversal experiments performed in this work represents the result of the conversion of a “disordered calcite” to aragonite on quenching. Reaction kinetics was found to be in the order of tens of hours at temperatures of 700 °C to achieve a complete transformation (Lin and Huang, 2004). This reconstructive transformation is therefore expected to be slow enough to be quenchable as drop in temperature from near solidus conditions to 700 °C occurs in a few seconds in multianvil apparatus.

Experimental brackets from this study, combined with data from previous works, were then used to constrain phase transition boundary from calcite V to aragonite, as well as CaCO_3 melting curve.

5. Thermodynamic modelling of Calcite V and CaCO_3 L properties

196 5.1 Locating the phase boundaries

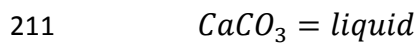
197 The transition boundary was thereby obtained by fitting the experimental brackets from this
198 study with the results from the ex-situ experimental results by Irving and Wyllie (1975), Buob
199 (2003) and Shatskiy et al (2014), an experimental dataset that does not support a positive change in
200 slope with increasing temperature as suggested by heating cycle experiments by Suito et al., (2001),
201 and by Litasov et al (2017)).

202 The calcite V-aragonite transition boundary can be expressed by the following first-order
203 equation:

$$204 \quad T_{ccv-arag} = 320.17 \times P + 397.6 \quad (1)$$

205 (P in GPa; T in K). This equation was then used, along with available thermodynamic data on
206 aragonite, to derive the thermodynamic properties of calcite V.

207 The melting curve for CaCO_3 was experimentally bracketed from 3-6 GPa. Melting of the
208 system was determined based on textural criterion through secondary electron imaging. Neither Ca-
209 oxides nor graphite nor gas bubbles were detected; run products were then interpreted in terms of
210 congruent melting:



212 since the quenched material consisted entirely of calcite. It is worth mentioning that incongruent
213 melting of CaCO_3 is expected to occur at low pressure (Wyllie and Boettcher, 1969).

214 CaCO_3 -melting curve was determined by quadratic least square regression of experimental
215 brackets from this study combined with previously published data, and finally expressed by the
216 following quadratic polynomial between 0 and 4.5 GPa:

$$217 \quad T_m = 1578.9 + 139.65 \times P - 11.646 \times P^2 \quad (2)$$

218 , where P is in GPa and T in K. Equation 2 was successively used for modelling the thermodynamic
219 properties for CaCO₃-liquid.

220 **5.2 Retrieving thermodynamic data for calcite V and liquid CaCO₃**

221 Thermodynamic data on the pure substances CaCO₃, MgCO₃, and FeCO₃ constitute the basis for
222 modelling a ternary carbonate melt solution model, which is crucial for predicting natural
223 phenomena involving carbonates stability, carbonatite segregation, and metasomatic processes in
224 the mantle. However, while some properties have been already derived for iron (Kang et al., 2015)
225 and magnesium (Kang et al., 2016) carbonates, a model for high-temperature calcite consistent with
226 liquid calcium carbonate is still lacking. It is beyond the goal of this work to assess a model offering
227 a complete description of order-disorder phenomena accounting for the complexities of the
228 transformation from calcite I to calcite IV, and ultimately to calcite V (Ishizawa et al., 2013). Our
229 model is intended to reproduce properties and phase relationships at mantle temperatures and
230 pressures where either aragonite or calcite V is expected to be the stable phase.

231 Here we derive the standard molar Gibbs free energies G_0 (J), entropies S_0 (J/K), and selected
232 volumetric properties along with a revision of heat capacities at high temperature, by fitting the
233 energy residue at conditions of the calcite V-aragonite transition boundary (Eq. 1), where $G_{ccv} =$
234 G_{arag} , and along the CaCO₃ melting curve (Eq. 2), where $G_{ccv} = G_{CaCO3L}$.

235 Provided the large uncertainties at high temperature and the lack of data for ccV, the
236 temperature dependence of thermal expansivity (α_T), and of bulk modulus (K_T), are defined by the
237 approximations of Holland and Powell (2011) as:

$$238 \quad \alpha_T = \alpha_0 \times \left(1 - \frac{10}{\sqrt{T}}\right) \quad (3)$$

$$239 \quad K_T = K_0 \times [1 - 1.5 \times 10^{-4} \times (T - 298.15)] \quad (4)$$

240 Volume at reference pressure $V(T, P_r)$ is

$$V(T, P_r) = V_0 \times [1 + \int_{T-T_r}^T \alpha(T, P_r) dT] \quad (5)$$

, while volumes at elevated pressures are computed by the Murnaghan equation of state:

$$V(T, P) = V(T, P_r) \times \left[1 - K' \times \frac{P}{(K' \times P + K(T, P_r))} \right]^{\frac{1}{K'}} \quad (6)$$

The isobaric heat capacity function $C_p(T)$ adopted is from Holland and Powell (1998):

$$C_p = a + b \times T + \frac{c}{T^2} + \frac{d}{\sqrt{T}} \quad (7)$$

Coefficients for heat capacity functions for aragonite and calcite V were modified to converge to the Dulong-Petit limit at high temperature (> 2000 K) as:

$$C_p = 3 \times R \times n + \alpha_T^2 \times V_T \times K_T \times T \quad (8)$$

where R is the gas constant and n the number of atoms in the substance of interest. At low temperatures (< 800 K), the C_p functions converge the heat capacities after Holland and Powell (2011).

The heat capacity function for liquid CaCO_3 has the same form of that used for solids, as discussed in detail in Kang et al. (2015).

Gibbs free energies of crystalline calcite V, and of CaCO_3L , at the P-T conditions of the calcite V-aragonite transition, given by Eq. 1, and the CaCO_3 melting reaction, given by Eq. 2, respectively, were calculated from:

$$\begin{aligned} G(P, T) &= G(P_r, T_r) - \int_{P_r, T_r}^{P, T} S(P_r, T) dT + \int_{P_r, T_r}^{P, T} V(P, T) dP \\ &= G(P_r, T_r) + \int_{P_r, T_r}^{P, T} C_p(P_r, T) dT - T * \int_{P_r, T_r}^{P, T} \left[\frac{C_p(P_r, T)}{T} \right] dT + \int_{P_r, T_r}^{P, T} V(P, T) dP \end{aligned} \quad (9)$$

The thermodynamic analysis was performed by adopting $S_0 = 87.99$ J/K of aragonite from Staveley and Linford (1969), later reported in Robie and Hemingway (1995), and G_0 of aragonite

261 averaging data from Robie and Hemingway (1995), and Königsberber et al. (1999). V_0 , α_0 , K_0 and
 262 K' of aragonite from the data presented in Palaich et al. (2016) re-fitted with eq. (6). The volume
 263 equation of state of calcite V is substantially unknown at high temperature; we therefore collected
 264 in-situ X-ray diffraction on CaCO_3 in order to determine thermal expansivity. Thermal expansion of
 265 calcite results increases from $4.1\text{E-}5 \text{ m}^\circ\text{C}$ determined in the 25-500 °C interval (Merlini et al.,
 266 2016) up to $1.34\text{E-}4 \text{ m}^\circ\text{C}$ in the highest temperature range investigated here. As a consequence G_0
 267 and S_0 of calcite V were fitted adopting V_0 at 3.60 J/bar, and fitting α_0 constrained at values higher
 268 than $1.0\text{E-}4 \text{ m}^\circ\text{C}$. Bulk modulus of calcite and its derivative were derived from Merlini et al
 269 (2016). Thermodynamic properties for liquid CaCO_3 were derived assuming a fixed bulk modulus
 270 derivative ($K' = 4.0$) (Kang et al., 2016). The results of the non-linear least-square fit are presented
 271 in Table 2.

272 The dependence of density on temperature and pressure for liquid CaCO_3 was calculated by
 273 Perple_X thermodynamic software (Connolly, 2005), at conditions of 1773 and 2073 K. Results are
 274 compared with those provided by molecular dynamics (MD) calculations implemented by empirical
 275 pair potential force field, and those evaluated by first-principle molecular dynamics (FPMD)
 276 proposed by Vuilleumier et al. (2014) fitted with Birch-Murnaghan equation of state and those
 277 obtained with the radial distribution functions by Hudspeth et al. (2018). The EoS retrieved from
 278 our study results in a compressibility for the CaCO_3 -liquid intermediate between those proposed by
 279 Vuilleumier et al. (2014) and Hudspeth et al. (2018) (Fig. 5). It is worth pointing out that our
 280 thermodynamic data retrieved for liquid CaCO_3 entirely derive from the melting curve of calcite V,
 281 with no additional constrain; as a consequence the agreement of the pressure dependence for density
 282 as calculated here and as obtained by independent methods is quite remarkable.

283 6. Discussion

284 6.1 Phase relationships in the system CaCO_3

285 Phase relationships have been calculated in the system CaCO_3 at pressures to 10 GPa using the
286 thermodynamic parameters for CaCO_3 provided from this study, and, for comparison, from Holland
287 and Powell (2011), and 2002 update of database in Holland and Powell (1998), hereafter
288 abbreviated as HP11 and HP02, respectively. The calculated results are compared with previously
289 published experimental data by Irving and Wyllie (1975) (IW75), Suito et al. (2001) (S01), Buob
290 (2003) (B03), Shatskiy et al. (2014) (S14) and Li et al. (2017) (Li17) (Fig.3).

291 As previously stated, it is beyond the goal of this work to offer a model reproducing the
292 transformations calcite I-IV-V. Thermodynamic data for calcite V were not optimized at low
293 temperature, resulting in a stability field for disordered calcite extending down to temperature of
294 approximately 610 °C; it is worth remembering that disordering to calcite IV in calcite was
295 observed from approximately 700 °C at nearly room pressure (Ishizawa et al., 2013)

296 At relatively low-pressures and low-temperature, the calcite V-aragonite boundary modelled
297 by employing our dataset does not differ significantly with that predicted by HP11, and HP02,
298 database (dashed, and dotted lines in Fig. 3, respectively). However, with increasing temperature,
299 larger deviations occur among the calculated results: HP11, and HP02 databases predict an
300 increasingly steepening slope for the calcite V - aragonite boundary, such that aragonite is not
301 expected to participate to melting in the uppermost mantle. On the contrary, the boundary
302 determined by our model maintains an almost constant dP/dT , then encountering the invariant point
303 calcite V - aragonite - liquid at about 5 GPa and 1650 °C, which matches fairly well with the
304 invariant point experimentally extrapolated by Irving and Wyllie (1975), and the shape of the
305 solidus, as determined by Li et al. (2017). Additionally, the stability field that we report for
306 aragonite covers the P-T conditions where aragonite was observed as the stable polymorph by Buob
307 (2003), and by Shatskiy et al. (2014).

308 At pressure above 8 GPa, the melting curve exhibits a moderately negative slope, then
309 suggesting that the liquid becomes more compressible than the solid, as it was also expected by
310 both *ab initio* simulations and *in-situ* conductivity experiments by Li et al. (2017).

311 **6.2 Modelling experimentally constrained carbonated-eclogite**

312 To further test the reliability of our thermodynamic model, pseudosections have been
313 performed on experimentally constrained complex chemical systems (Table 3), such as *bulk*
314 compositions OTBC (carbonated eclogite with the altered basalt composition (OTB) + 10.1 wt%
315 calcite + 0.12 wt% water bulk composition, Hammouda, 2003), SLEC1 (carbonated eclogite,
316 Dasgupta et al., 2004), GA1cc (eclogite composition of GA1, representing altered oceanic MOR
317 basalt, +10% CaCO₃, Kiseeva et al., 2013) and EC1 (carbonate-bearing eclogite assemblages,
318 Yaxley and Brey, 2004), that are generally referred to as mantle heterogeneities derived from deeply
319 recycled, subducted oceanic crust (Fig. 6 and 7). For comparison, phase relationships have been
320 modelled by using the thermodynamic properties for CaCO₃ from both this study, and HP11
321 database.

322 The bulk composition OTBC (Hammouda, 2003) represents a partially hydrated carbonated
323 eclogite (0.12 wt.% H₂O in the starting material). Its re-equilibration at mantle conditions, promoted
324 by the deep recycling of oceanic material, may account for the extraction of carbonate-rich liquids.
325 We performed pseudosections with the Perple_X thermodynamic calculators (Connolly, 2005), on
326 a K-free OTBC from 1-10 GPa and 900-1300 °C by implementing the thermodynamic dataset for
327 CaCO₃ retrieved from this study (Fig. 6a), and then compared the results with those obtained by
328 employing the parameters for CaCO₃ provided by the HP11 database (Fig. 6b). Solution models for
329 clinopyroxene, spinel, plagioclase, and garnet used here are from Jennings and Holland (2015), for
330 binary H₂O-CO₂ fluid from Connolly and Trommsdorff (1991), and for ternary Ca-Fe-Mg
331 carbonates from Franzolin et al. (2011). The latter model is able to predict immiscibilities in
332 carbonate solid solutions. The pseudosection presents a clinopyroxene - garnet - coesite assemblage

stable over a large range of P-T conditions (Fig. 6), while plagioclase only occurs at pressures lower than ~ 2.5 GPa. From 1 to ~ 4 GPa, the eclogitic assemblages coexist with a magnesian-calcite solid solution ($X_{\text{Ca}} > 0.8$), whereas at higher pressures the stable carbonate phases are aragonite, and aragonite + magnesite with increasing pressure. Aragonite stability field covers a wide P-T range, extending down to 4 GPa at 1300 °C; the abundance of aragonite calculated for this bulk composition reaches approx. 11 wt.% in the region 4 - 5 GPa at 1000 °C, and 7 - 10 GPa at 1300 °C.

The calculation performed with our dataset was found to well reproduce the experimental results especially in the high-pressure region, as the association aragonite + magnesite was detected at near-*solidus* conditions from 7 to 10 GPa (Hammouda, 2003). The abundance that have been obtained for aragonite agrees fairly well with its ubiquitous presence in the experiments at pressures higher than 6 GPa; above this pressure, aragonite was found to persist largely above the experimentally determined solidus (Fig. 6a), up to 1200 °C where it coexists with a carbonatitic melt (Hammouda, 2003). Although a model for carbonatitic liquids cannot be constructed here because of the large non-ideality of CaCO_3 - MgCO_3 - FeCO_3 plane, we expect that the calculated abundance of aragonite correlates with likelihood of its occurrence at supersolidus conditions, notably when the liquid is silicate. The development of a thermodynamic model for describing the liquidus surface of ternary carbonatitic liquids will be discussed elsewhere.

The phase diagram calculated by employing the CaCO_3 parameters from HP11 (Fig. 6b) reveals major differences in a P-T region, between 1100-1300 °C and 4-6 GPa, which is most relevant for magma generation and where transition from carbonate to silicate melt was inferred to occur (Hammouda, 2003). We speculate that aragonite is expected to be entirely consumed approaching the solidus according to HP11 database, whereas the adoption of our new thermodynamic dataset (Fig. 6b) show that aragonite might persist with the liquid, as observed at higher pressures. The implications of this will be discussed below.

358 The stability of aragonite within an eclogitic assemblages was also verified at conditions of the
359 mantle transition zone, by Kiseeva et al. (2013) on carbonated eclogite, GA1cc. Thermodynamic
360 modelling (not shown) define a dominant eclogitic association, given by clinopyroxene, garnet and
361 coesite/stishovite, coexisting with aragonite + magnesite.

362 In order to constrain the variability of aragonite occurrence within eclogitic systems where
363 CaCO_3 -saturation was not experimentally observed, pseudosections were also performed, by
364 employing our new dataset, on *bulk* compositions SLEC1 (Dasgupta et al., 2004) and EC1 (Yaxley
365 and Brey, 2004). The thermodynamic calculation performed on SLEC1 (K-free, Mn-free for sake of
366 simplicity) exhibits very small fractions of aragonite (<1 wt%), limited to pressures higher than 5
367 GPa (Fig. 7a); the computed results are in substantial accord with the experimental record, where
368 only calcian-dolomite solid solution, and magnesite at higher pressure, were found at near-*solidus*
369 conditions, whereas aragonite is not observed. Differently, the pseudosection calculation
370 performed on EC1 composition displays a large P-T range where aragonite forms, reaching modal
371 contents of up to 20 wt % at high pressure (Fig. 7b). However, aragonite was not observed by
372 Yaxley and Brey (2004), as the range of pressures and temperatures investigated (2.5-5.5 GPa and
373 ~1125-1310°C) were just below the stability field predicted by our calculations. The bulk
374 composition EC1 is most suitable to highlight the relevance of aragonite at upper mantle conditions.

375 **6.3 Aragonite stability in mantle heterogeneities**

376 The new set of thermodynamic parameters for CaCO_3 was found to describe properly phase
377 relationships in carbonated eclogites, which are regarded as potential sources for feeding
378 carbonatitic and silica undersaturated basaltic magmas observed within intra-plate environments
379 (Sobolev et al., 2005; Aulbach et al., 2017). The geochemical signature exhibited by these
380 magmatic suites are interpreted to derive from an eclogitic component, inherited by a deeply
381 subducted oceanic crust (Hofmann, 1997; Kogiso et al., 1997); therefore, the *bulk* composition of
382 such recycled material may span a wide compositional range in major elements (Hirschmann and

383 Stolper, 1996), notably due to the heterogeneous extent of the primeval ocean floor alteration, and
384 of the subsequent dehydration processes during subduction. Phase assemblages and melting
385 reactions within mantle heterogeneities are not univocal in the P-T space, but largely differ
386 depending on several compositional effects.

387 The initial *bulk* #Ca ratio, $\text{Ca}/(\text{Ca} + \text{Mg} + \text{Fe})$ was found to strongly determine the carbonated-
388 eclogite melting temperature and chemical features of liquid produced, due to the location of the
389 minimum of the liquidus surface on the binary $\text{CaCO}_3\text{-MgCO}_3$ (Irving and Wyllie 1975). The
390 composition of dolomite, and magnesian-calcite solid solution at near-*solidus* is buffered by the
391 partitioning of Ca-Mg with silicates, mainly garnet and clinopyroxene, which in turn depends on the
392 contents of Al_2O_3 and Na_2O in the bulk composition. These compositional parameters partly
393 account for the large discrepancies on the location of the carbonated-eclogite *solidus* curve, which
394 may differ by the order of ~ 200 °C between the experimental determination on SLEC1 (Dasgupta
395 et al., 2004), EC1 (Yaxley and Brey, 2004), and OTBC (Hammouda, 2003). The very high melting
396 temperature recorded on EC1 is likely to be related to the high #Ca values (#Ca = 0.435), along
397 with high CO_2 contents (15.38 wt.%), with respect to the SLEC1 composition (#Ca = 0.319, CO_2 =
398 5 wt.%), whereas the relatively low-T *solidus* determined on OTBC is probably due to the presence
399 of H_2O , which is known to strongly depress the melting temperature on #Ca-rich systems (Poli,
400 2015).

401 Aragonite is often referred to as a phase relevant at relatively low-T conditions, and commonly
402 associated to subduction zone processes. However, the relevance of aragonite in the generation and
403 evolution of igneous suites is testified by its occurrence in metasomatized pyroxenites from the
404 North Andean mantle (Ferri et al, 2017) and in alkaline lavas from Clatrava, Spain (Humphreys et
405 al., 2010). The stability of aragonite not simply correlates to the amount of CaO in the bulk
406 composition, but more closely relates to availability of alumina. The higher the Al_2O_3 , the larger the
407 fraction of Mg and Fe fractionated in garnet, the higher the “residue” of CaO available to form a

408 calcium carbonate component (Fig. 8), if CO₂ is available. As a consequence, although the *bulk*
409 composition SLEC1 adopted by Dasgupta et al. (2004) to model the solidus of carbonated eclogites
410 is relatively enriched in CaO, the low alumina content drives the composition toward the
411 (Mg,Fe)CO₃ component, when compared to other experimentally investigated *bulk* compositions
412 (compare Figs. 6 and 7).

413 The role of aragonite on controlling the melting temperatures in eclogitic systems remains
414 unclear, but the relevance of the assemblage aragonite + magnesite-siderite solid solution at high-
415 pressure, at near-*solidus* conditions, was both thermodynamically predicted (in this study) and
416 experimentally testified in Hammouda (2003), and Kiseeva et al. (2013). It is worth remembering
417 that a carbonate, namely aragonite, and graphite or diamond are not mutually exclusive in complex
418 bulk composition. A field exists in the log *f*O₂ vs. P-T space where carbonates coexist with
419 elemental carbon, and the lower the temperature, the larger the log *f*O₂ interval (see Fig 3 in Poli et
420 al., 2009).

421

422 **7. Implications for the geochemical signature of anorogenic magmas**

423 The thermodynamic properties for aragonite, calcite V, and CaCO₃L, were derived here by a
424 least-square fitting analysis of the experimental data available. The boundary of phase transition
425 from calcite V to aragonite, and the melting curve of CaCO₃, were provided in a P-T region relevant
426 for investigating processes in the upper mantle. We revisited the aragonite stability field, which was
427 found to extend to lower pressures and, relatively, higher temperatures in comparison to what it has
428 been recently assessed on the basis of in-situ experiments and previous thermodynamic modelling.
429 *In-situ* XRD studies (Suito et al., 2001; Litasov et al., 2017) restricted aragonite stability, yielding
430 results conflicting with *synthesis* experiments where aragonite has been observed up to the melting
431 curve at pressures as low as 4.7 GPa.

432 The modelling of carbonated-eclogites shows that aragonite may occur as a major compound
433 in a range of P-T conditions where melting reactions have been experimentally determined
434 (Hammouda, 2003; Kiseeva et al., 2013). It is widely accepted since long time that the composition
435 of carbonatitic and kimberlitic melts requires a metasomatized mantle source (Wyllie, 1980;
436 Ringwood et al., 1992; Eggler, 1978). The extended stability of aragonite may drive significant
437 fractionation of trace elements in such coexisting liquids because solid-melt trace elements partition
438 coefficients for aragonite are drastically different from those determined for magnesite or silicates
439 (Brey et al., 2015). Aragonite strongly sequesters REE, notably intermediate REE, and Sr, whereas
440 HFS elements result highly incompatible, to a larger extent than magnesite (see Fig. 8 in Brey et al.,
441 2015). Figure 9 highlights the peculiar geochemical signature of aragonite, with reference to the
442 Primitive Mantle of Sun and McDonough (1989). Fractionation of aragonite from a segregated,
443 mobile, volatile enriched liquid would therefore contribute to the definition of a typical anorogenic
444 signature as observed for those silica undersaturated magmas, where a carbonatitic component is
445 claimed, including kimberlites (Becker and Le Roex, 2006).

446 As H₂O strongly depresses the *solidus* temperature of carbonated lithologies enriched in Ca,
447 the generation of hydrous carbonatitic liquids is potentially extended to subduction zone tectonic
448 settings (Poli 2015, 2016). How aragonite participates to melting reactions at such conditions is
449 currently unknown, and trace element pattern fractionation might significantly differ from
450 anorogenic settings, provided the expected dissimilarity in H₂O availability and thermal field.

451 Aragonite is expected to be a major player in the evolution of a mantle refertilized by deeply
452 subducted altered oceanic crust, as it persists to hot mantle adiabats and provides a mean to
453 fractionate trace elements at the onset of melting. Further investigations are mandatory for
454 unravelling the complex interplay between carbonates and H₂O components in mantle
455 heterogeneities, and how their relationships eventually govern the geochemical evolution of intra-
456 plate magmatism.

457

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466

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667 **Figure captions**

668 **Figure 1**

669 Summary of the phase relations in the system CaCO_3 . Phase boundaries from calcite I (CCI) to
670 calcite IV (CCIV), and calcite V, are labelled as M76-Mirwald (1976), and BS03-Bagdassarov and
671 Slutskii (2003). Transition curves from calcite to aragonite as experimentally investigated by L17-
672 Litasov et al. (2017), S01 - Suito et al. (2001), M76 - Mirwald (1976), and IW75-Irving and Wyllie
673 (1975), predicted by calorimetric measurements by R89-Redfern et al. (1989), and calculated with
674 thermodynamic database from HP11-Holland and Powell (2011), and HP02-Holland and Powell
675 (1998). Open diamonds represent experiments at 6 GPa by B03 - Buob (2003) where aragonite was
676 observed as the stable CaCO_3 -polymorph. Open triangle and filled triangle represent experiments
677 by S14 - Shatskiy et al. (2014) where aragonite and melt, respectively, have been detected. Melting
678 was investigated by S18-Shatskiy et al. (2018), Li17-Li et al. (2017), IW75-Irving and Wyllie
679 (1975), and S01-Suito et al. (2001). The sub-oceanic ridge and ocean islands geotherms are from
680 Dasgupta and Hirschmann, 2010.

681

682 **Figure 2**

683 Secondary electron image of fibrous crystals of calcium carbonate (Run products CCMS1: 4.7 GPa-
684 1700 °C). These morphologies are interpreted as disequilibrium textures developed during rapid
685 cooling from a CaCO_3 -liquid present at run conditions.

686

687 **Figure 3**

688 Experimental results and phase boundaries for the system CaCO_3 , as predicted by employing
689 thermodynamic parameters from this study (solid line), HP11 database (dashed line), and HP02

690 update (dotted line). Results are compared with experimental data from IW75 (Irving and Wyllie,
691 1975), Li17 (Li et al., 2017) with “Pt sphere” (falling sphere) and ECM (electrical conductivity
692 measurements) experiments, B03 (Buob, 2003), S14 (Shatskiy et al., 2014), and S01 (Suito et al.,
693 2001). Filled green, open red, and filled violet symbols represent experimentally detected aragonite,
694 calcite V, and melt, respectively.

695

696 **Figure 4**

697 Unit cell volume of CaCO_3 vs. temperature data from this study at ambient pressure. The filled
698 circles represent the phase calcite - I. The open circles are phase calcite - IV and filled triangles are
699 disordered phase of calcite - V.

700

701 **Figure 5**

702 Isothermal compression curves for CaCO_3 at 1773 K (green) and 2073 K (red), determined from
703 this study (solid lines), and by MD data fitted with Birch-Murnaghan equation of state by
704 Vuilleumier et al. (2014) (dashed lines). The red/green dots with pressure error bars are obtained by
705 FPMD calculations (Vuilleumier et al., 2014) and the red/green dots with pressure and density error
706 bars are obtained with the radial distribution functions by Hudspeth et al., (2018).

707

708 **Figure 6**

709 Calculated pseudosections for carbonated eclogite *bulk* composition of OTBC, modified (K_2O -free)
710 after Hammouda (2003). The thermodynamic parameters for CaCO_3 are provided by: (a), this
711 study; (b), Holland and Powell (2011). Solid solution models here used are Cpx(JH), Sp(JH),
712 Pl(JH), and Grt(JH) proposed by Jennings and Holland (2015), F by Connolly and Trommsdorff

713 (1991), and oCcM(EF) (Mg-Fe-Ca carbonate solid solution) by Franzolin et al. (2011). Carbonatitic
 714 *solidus* (purple curve), and silicate *solidus* (green curve), were experimentally determined by
 715 Hammouda (2003). Crosses show all the experiments with aragonite from Hammouda (2003), and
 716 the diamond shows the experiment with aragonite in this P-T region from Kiseeva et al., (2013).
 717 Red contours represent calculated modal contents of aragonite (wt.%). Phase abbreviations are: cpx
 718 - clinopyroxene, pl - plagioclase, grt - garnet, ky - kyanite, sp - spinel, F - fluid, mg-cc - magnesio-
 719 calcite_{s.s.}, mst - magnesite, arag - aragonite, q - quartz, coe - coesite, stv - stishovite.

720

721 **Figure 7**

722 Calculated pseudosections for carbonated eclogite *bulk* composition of a) SLEC1 (Dasgupta et al.,
 723 2004), and b) EC1 (Yaxley and Brey, 2004), by employing the thermodynamic parameters for
 724 CaCO₃ retrieved from this study. *Bulk* compositions have been simplified by neglecting the TiO₂,
 725 MnO, Cr₂O₃, and K₂O components. Green *solidus*, and blue *solidus* curves are provided by
 726 Dasgupta et al. (2004), and Yaxley and Brey (2004), respectively. Red contours represent calculated
 727 modal contents of aragonite (wt.%). Phase abbreviations are: cpx - clinopyroxene, ol - olivine, grt -
 728 garnet, sp - spinel, mg-cc - magnesio-calcite_{s.s.}, cc-dol - calcio-dolomite_{s.s.}, mst - magnesite, arag -
 729 aragonite.

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732 **Figure 8**

733 Projected *bulk* compositions of reference carbonated eclogites, in a ternary CaCO₃-MgCO₃-
 734 NaAlSi₂O₆ compositional space: OTBC (Hammouda, 2003), SLEC1 (Dasgupta et al., 2004),
 735 GA1cc and Volga (Kiseeva et al., 2013), and EC1 (Yaxley and Brey, 2004). *Bulk* compositions

736 were projected from $\text{SiO}_2 + \text{TiO}_2 + \text{CO}_2 + \text{Fe} \leftrightarrow \text{Mg}$ (exchange vector) + ternary garnet
737 $(\text{Mg}_{0.5}\text{Fe}_{1.5}\text{Ca}_{1.0})\text{Al}_2\text{Si}_3\text{O}_{12}$.

738

739 **Figure 9**

740 $[\text{La}/\text{Nb}]_{\text{norm.}}$ versus $[\text{La}/\text{Sm}]_{\text{norm.}}$ diagram, normalized to the Primitive Mantle (PM) of Sun and
741 McDonough (1989). Coloured diamonds refer to experimentally synthesized phases on carbonate
742 sediments at 7.5 GPa - 900 °C (Brey et al., 2015): violet - aragonite, red - melt, blue - magnesite,
743 green - garnet. Red, and orange circles represent Group I, and Group II close-to-primary kimberlite
744 compositions, respectively (Becker and Le Roex, 2006). OIB and N-MORB compositions are from
745 Sun and McDonough (1989), primitive arc andesites composition from Kelemen et al. (2003). A
746 schematic representation is reported for coexisting phase assemblage in the experiments of Brey et
747 al. (2015), on the basis of topologies discussed in Poli (2016).

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753 **Table 1**

754 Experimental run conditions and run products in the pure CaCO₃ system. M - melt, Cc - calcite, Ara
755 - aragonite.

Run	P (GPa)	T (°C)	Run time (hours)	Run products
CCMS1	4.7	1700	1.55	M
CCMS3	3.0	1650	2.6	M
CCMS4	3.8	1700	0.83	M
CCMS18	3.8	1500	24.17	Cc
CCMS19	4.7	1500	22.3	Ara
CCMS20	4.3	1500	17.5	Cc + Ara
CCMS21	3.3	1300	20	Cc
CCMS28	3.8	1300	27.42	Cc
CCMS30	4.2	1300	24.23	Ara
CCMS31	4.7	1650	2.0	Ara
CCMS34	6.0	1750	2.0	M

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764 **Table 2**

765 Standard molar thermodynamic properties of Aragonite, Calcite V and CaCO₃L at 298.15 K and 1
766 bar. G₀ - Gibbs free energy, H₀ - enthalpy, S₀ - entropy, V₀ - volume, C_p - heat capacity,
767 α₀ - thermal expansivity, K₀ - bulk modulus, K' - pressure derivative of the bulk modulus.

	G ₀ (J)	H ₀ (J)	S ₀ (J/K)	V ₀ (J/bar)	C _{p,298.15} (J/K)	α ₀ (1/K)	K ₀ (bar)	K'
Aragonite	-1233820	-1207586	87.99	3.395	81.2	1.082E-4	689960	4.397
Calcite V	-1229812	-1199884	100.38	3.60	83.3	1.10E-4	725200	4.0
CaCO₃L	-1174842	-1139660	118.0	4.045	125.7	2.473E-4	78000	4.0
Fitted coefficients of C _p referred to the polynomial Eq. (7)								
	a	b	c	d				
Aragonite	137.906	0.023	-602802.8	-980.885				
Calcite V	88.61	0.03945	-1094483.95	-81.726				
CaCO₃L	117.071	0.01409	-12434.329	79.396				

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776 **Table 3**

777 *Bulk compositions of reference carbonated eclogites expressed in wt.% oxides. OTBC, Hammouda*
 778 *(2003), SLEC1, Dasgupta et al. (2004), EC1, Yaxley and Brey, (2004), GA1cc, Kiseeva et al.*
 779 *(2013). #Mg is Mg/(Mg+Fe_T) molar ratio. #Ca is Ca/(Ca+Mg+Fe_T) molar ratio.*

	OTBC	SLEC1	EC1	GA1cc
SiO ₂	47.23	41.21	30.11	45.32
TiO ₂	-	2.16	-	1.34
Al ₂ O ₃	15.35	10.89	11.74	14.88
Cr ₂ O ₃	-	0.09	-	-
FeO _T	8.93	12.83	10.05	8.85
MnO	-	0.12	-	0.15
MgO	6.24	12.87	12.44	7.15
CaO	14.77	13.09	19.41	14.24
Na ₂ O	2.91	1.63	0.87	3.14
K ₂ O	0.02	0.11	-	0.40
P ₂ O ₅	-	-	-	0.14
CO ₂	4.43	5.00	15.38	4.40
H ₂ O	0.12	-	-	-
Total	100.00	100.00	100.00	100.00
#Mg	0.555	0.641	0.688	0.590
#Ca	0.485	0.319	0.435	0.458

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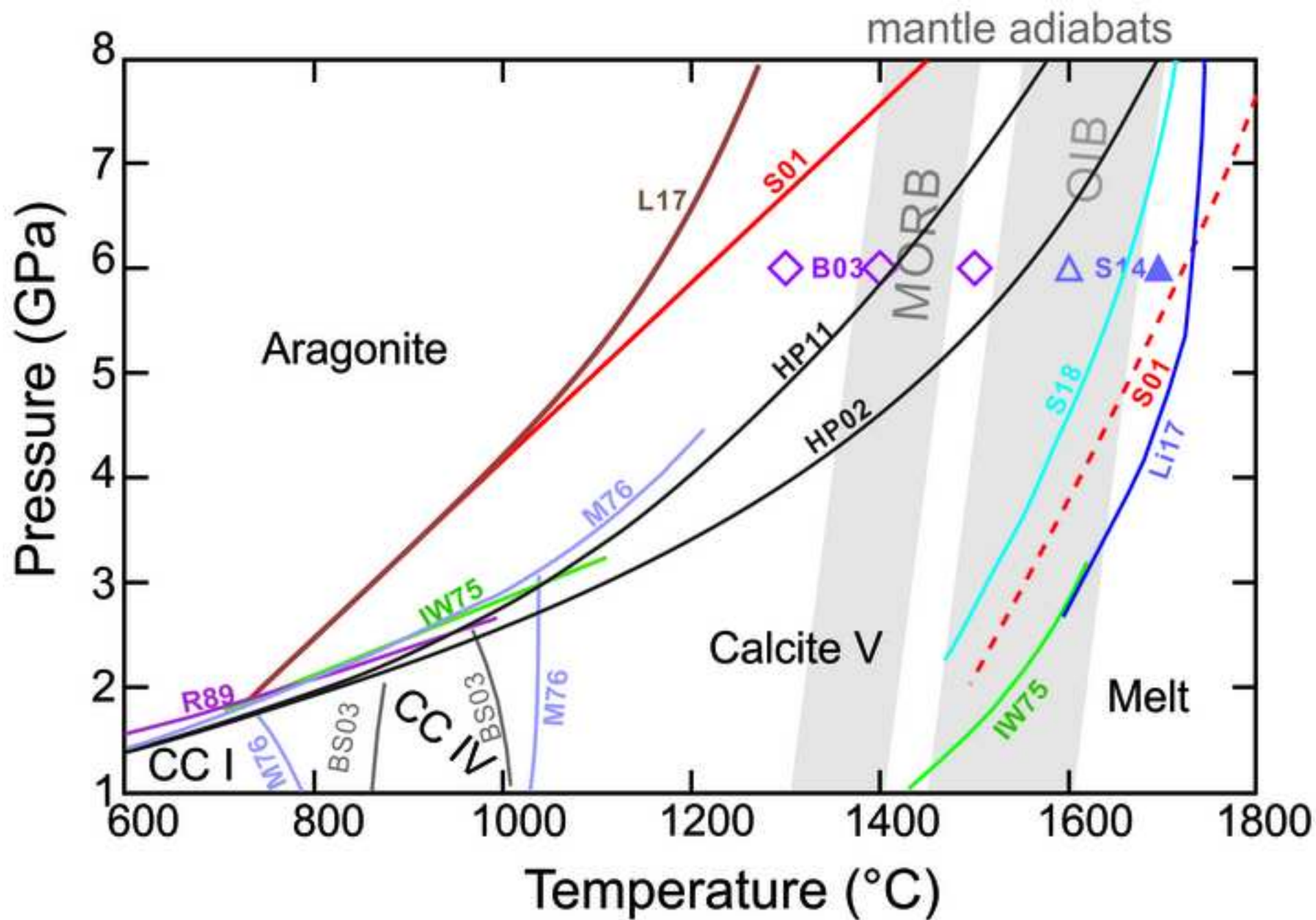


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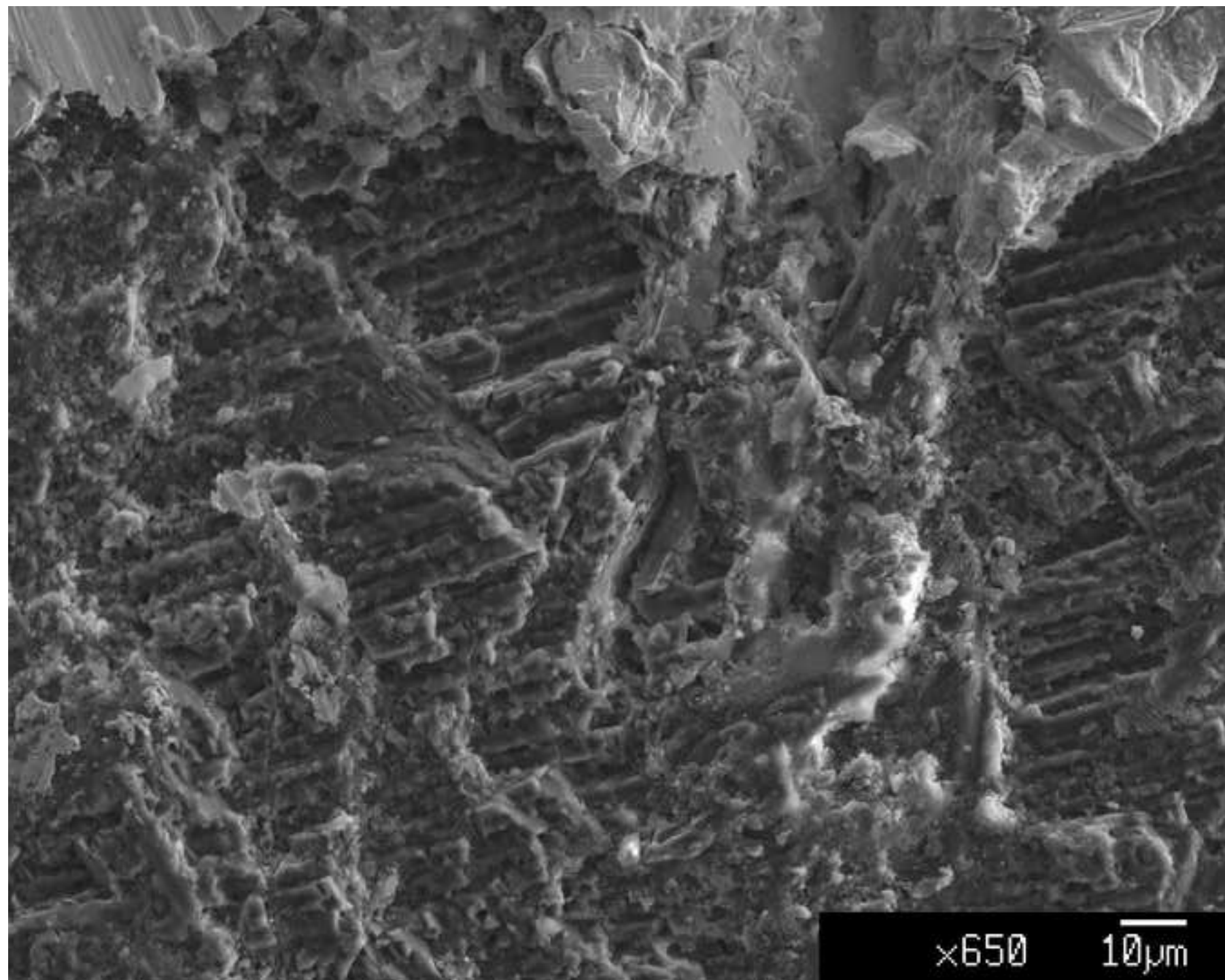


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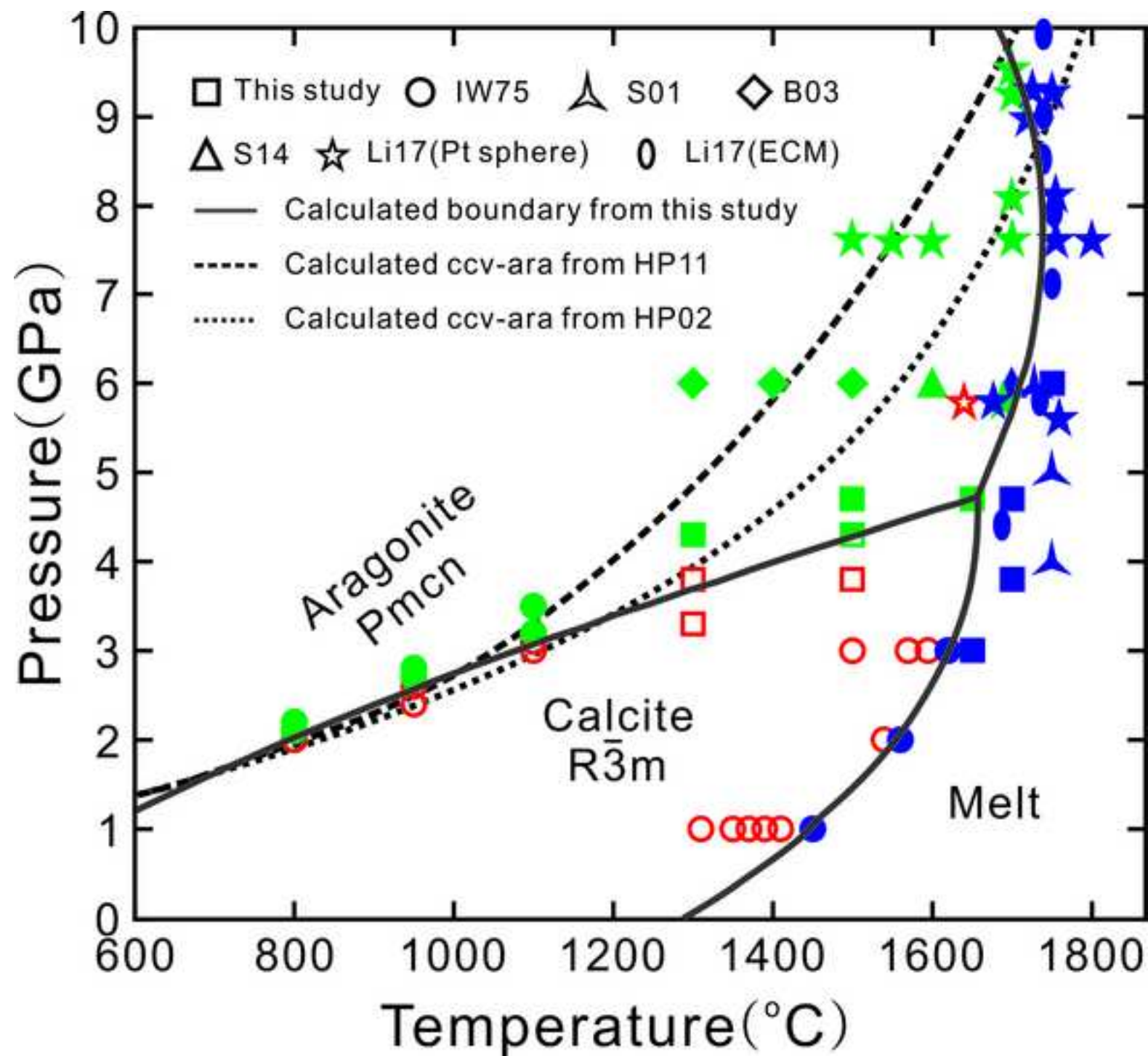


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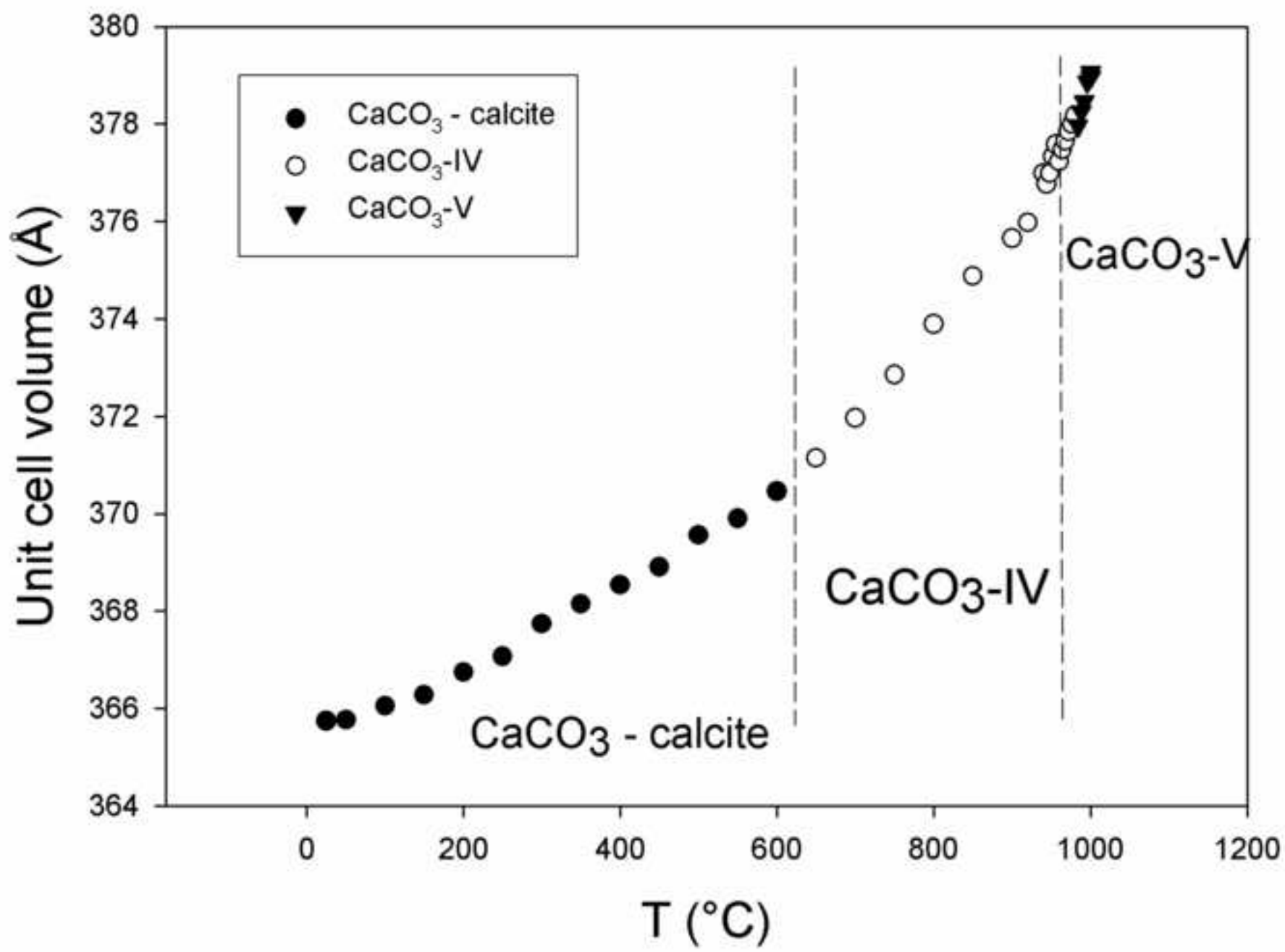


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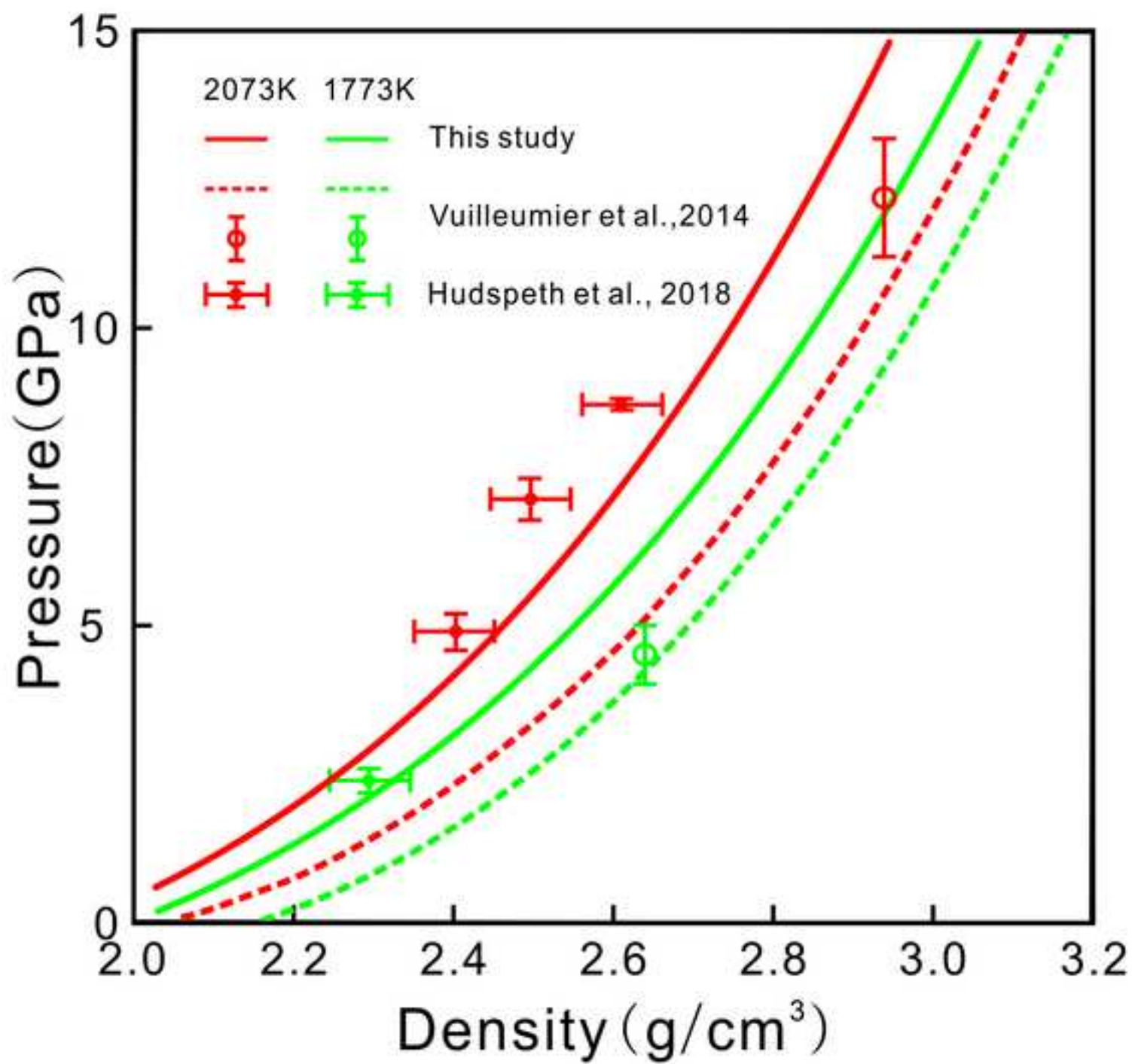


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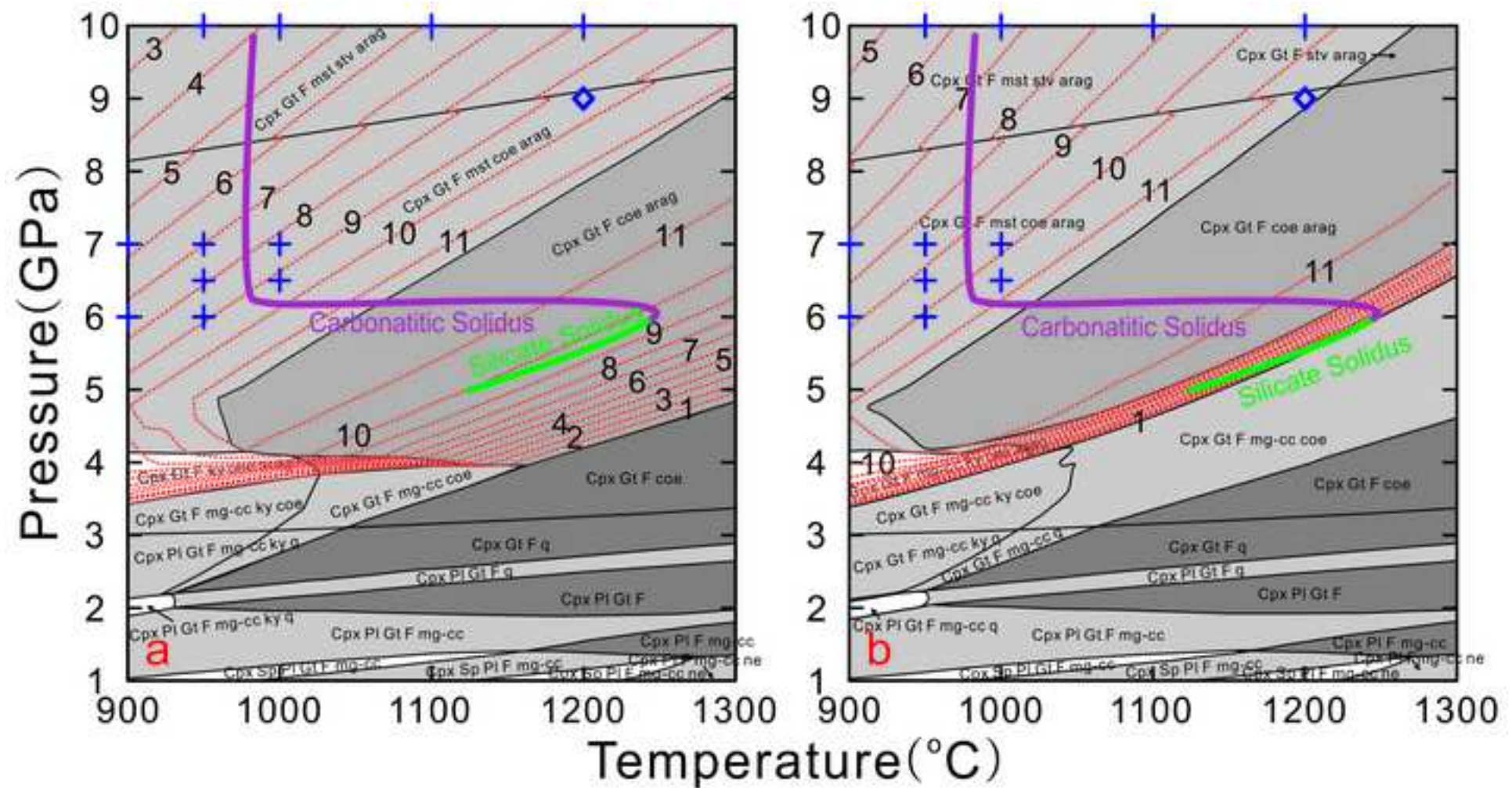


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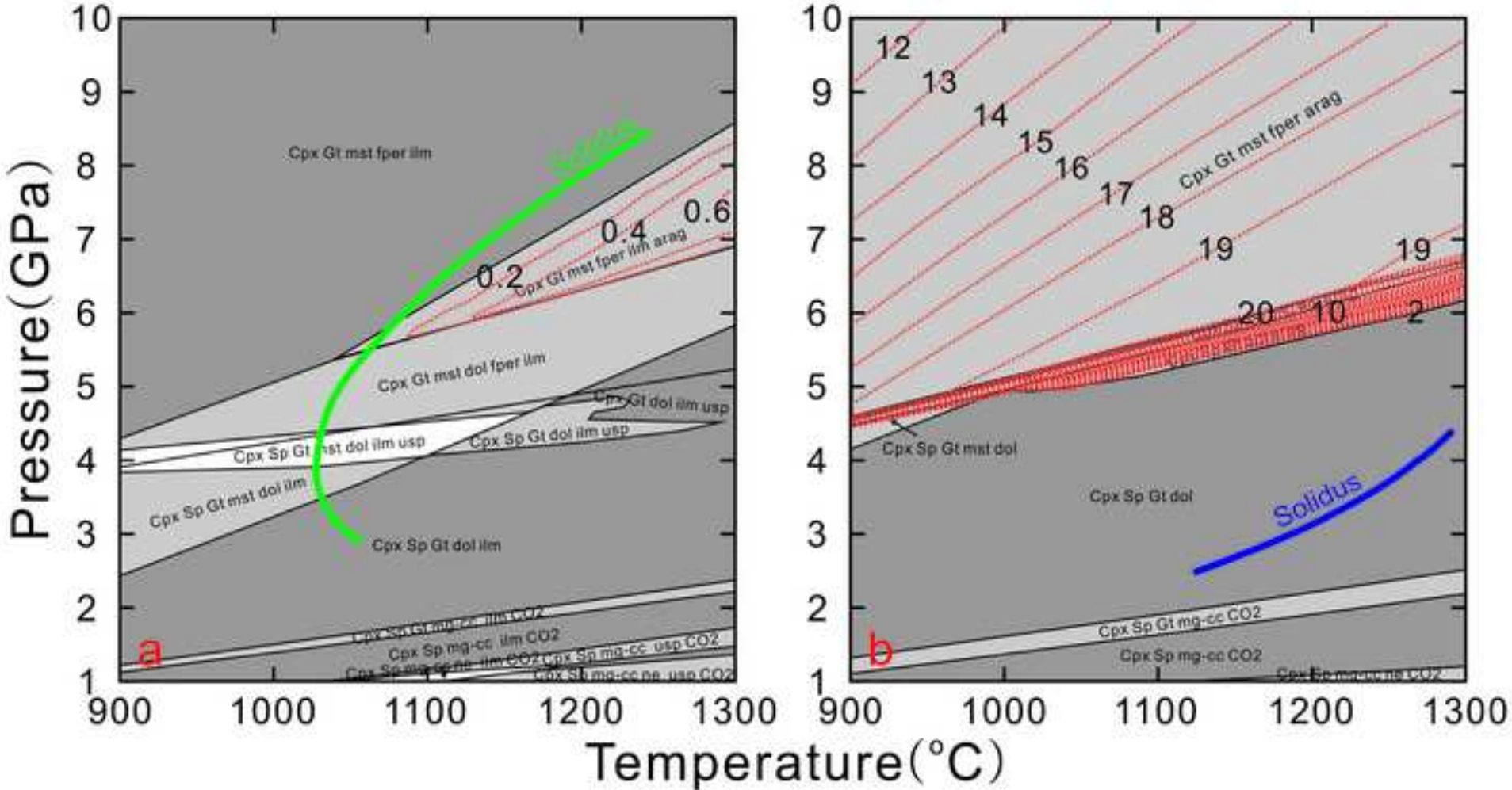


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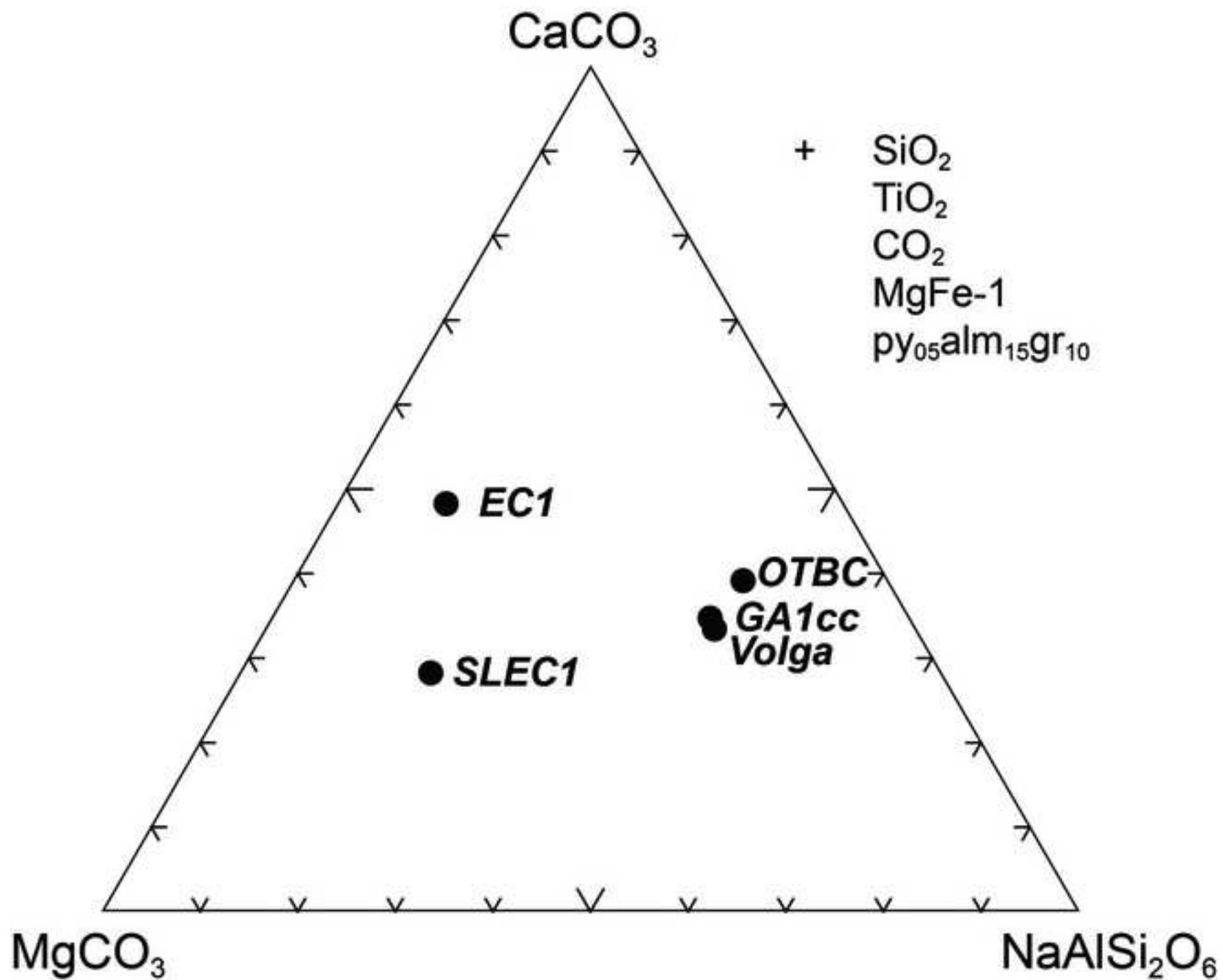
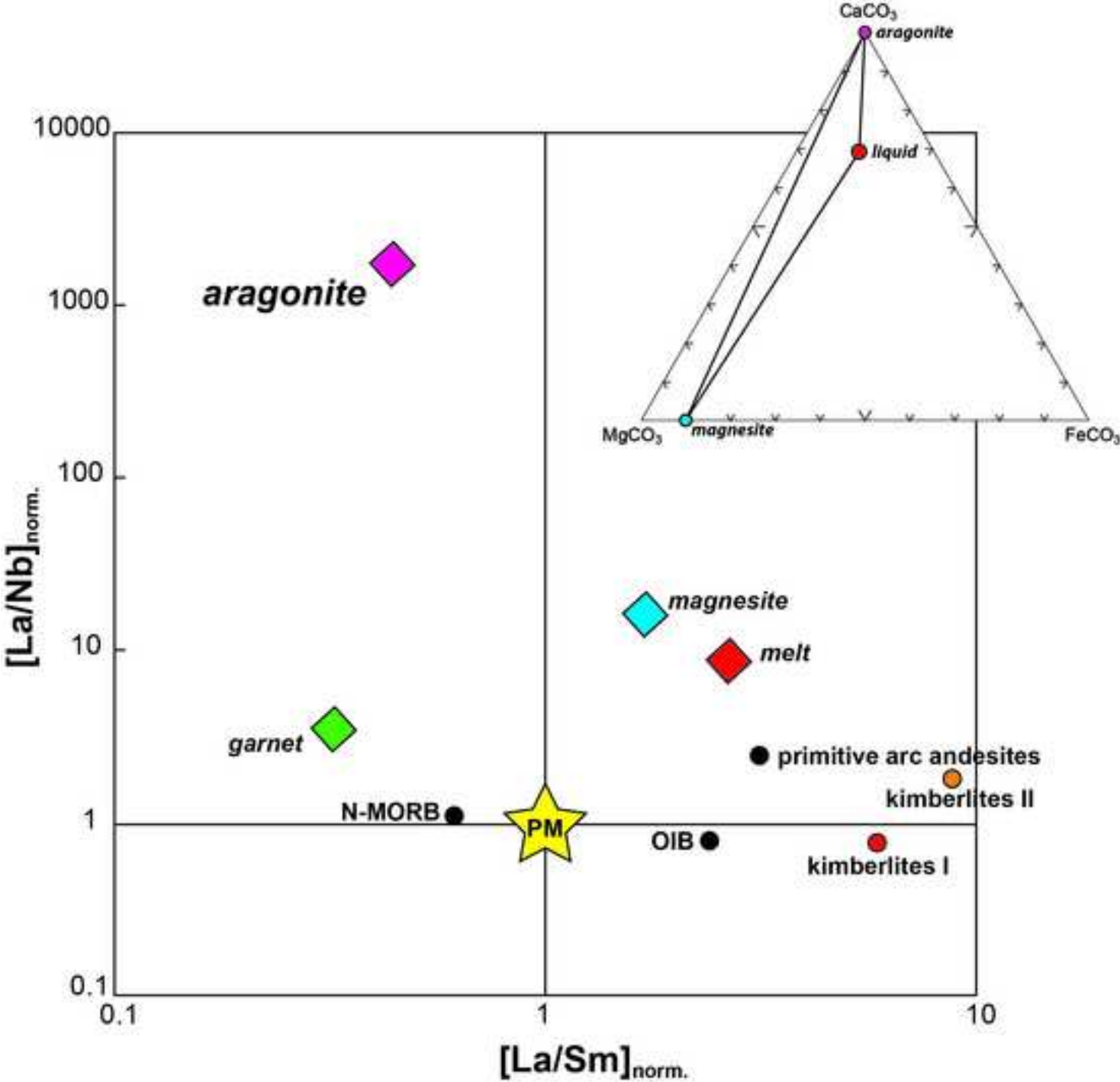


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